Cryptorchidism And Hyperthermia Induced Testicular Injury In Adult Male Rats; Intervention Of Alpha Tocopherol

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ABSTRACT

Introduction: The objectives of this study is to determine the effects of cryptorchidism and hyperthermia on the testicular cytoarchitecture of adult male rats, quality and quantity of semen and the level of oxidative stress in the
Methods: Twenty four (24) adult Wistar rats were used for this experiment and they were divided into seven groups (n=4). Experimental cryptorchidism was carried out in some animals while others were subjected to hyperthermia from two sources (hot oven and light bulb) for 28 days; followed by intervention of 100mg/kg alpha-tocopherol for fourteen (14) days. All animals were then euthanized under anaesthesia. The testes were excised and fixed and processed using haematoxylin and eosin technique. Plasma was assayed for glutathione, superoxide dismutase (SOD) and malondialdehyde (MDA).

Discussion: Cryptorchidism and hyperthermia caused in cycto-architectural distortion in the testicular structures of exposed animals and significant reductions in caudal epididymal sperm count and motility, morphology and life and death ratio. Plasma SOD and GLU levels also reduced significantly in exposed animals while MDA levels increased significantly, indicating peroxidation of testicular tissue. Alpha-tocopherol due to its anti-oxidative properties impeded lipoperoxidation of the membrane, conserving cell union and increasing the amount of germ cells in the seminiferous epithelium.

Conclusion: At doses and durations tested, alpha-tocopherol (aT) reverted to a significant extent the effects of cryptorchidism and hyperthermia by reducing apoptosis in germ cells, promoting cell survival and reducing histological alterations to the seminiferous epithelium.

Keywords: Cryptorchidism, hyperthermia and Alpha-tocopherol.

INTRODUCTION
The testes function in production of the male gametes or spermatozoa and male sexual hormone, (hence referred to as double glands) which stimulates the accessory male sexual organs and causes the development of the male sex characteristics (Mescher, 2010; Lutz, 2009). In most mammals, the temperature around the testicular region is lower than the core body temperature. In man, scrotal temperature is 2-30C lower than rectal temperature and the optimum temperature. Impaired testicular thermoregulation is commonly implicated in abnormal spermatogenesis and impaired sperm function in animals and humans, with outcomes ranging from subclinical infertility to sterility (Thundathi et al., 2012). Oyewopo and Togun in 2005 reported that a small increase in the temperature of the testis does not destroy the germinal epithelium; however, it reduces testis weight and sperm production and brings a greater incidence of morphologically abnormal spermatids and spermatozoa (Oyewopo and Togun, 2005; Bedford, 1991). Occupational exposure to high temperatures adversely affects testicular function causing partial or complete spermatogenic arrest, also causes deterioration in sperm morphology and impairs motility. This leads to oligoasthenoterozoospermia (OAT) and azoospermia (Dada et al., 2001).

Cryptorchidism (undescended testes); the absence of one or both testes from the scrotum happens to be the most common birth defect of the male genitalia, which if persists could result in reduced fertility and increased risk of testicular
germ cell tumors (Wood and Elder, 2009). Cryptorchidism has been implicated to cause oxidative stress in animals that the process was performed on, it was reported that Cryptorchid rats had lower testicular weight, sperm count, germ cell count, testicular superoxide dismutase (SOD) concentration, testicular total protein and higher testicular malondialdehyde (MDA) concentration compared to animals in control group (Ayobami et al., 2013; Dutta et al., 2013). Alpha-Tocopherol (αT) is an important antioxidant that localizes to cell membranes. The primary function of vitamin E is as an antioxidant; because it helps reduce oxidation of lipid membranes and the unsaturated fatty acids and prevents the breakdown of other nutrients by oxygen. This protective, nutritional antioxidant function is also performed and enhanced by other antioxidants, such as vitamin C, beta-carotene, glutathione (L-cysteine), and the mineral selenium (Laila and Sahar 2009; Aybek et al., 2008; Traber and Kayden 1987). A significant reduction of lipoperoxidation was observed in the cryptorchid group treated with α-Tocopherol compared to the untreated cryptorchid group, at long term, an increase in the area and maturation of the seminiferous epithelium, a decrease in apoptosis and histological alterations and an increase in fertility was observed in animals treated with alpha-Tocopherol (Saalu et al., 2014; Rosa et al., 2011).

The broad aim of this study investigate the protective properties of alpha tocopherol (vitamin E) on cryptorchidic and heat (hyperthermia) induced testicular injury.

MATERIALS AND METHODS
The experiment was performed in conformity with the Rules and Guidelines of the Animal Ethics Committee of University of Ilorin. The experiments were conducted at the Department of Anatomy, College of Health Science, University of Ilorin.

Animals
Twenty four (24) male Wistar rats (150-210g) were purchased from Banky livestock company, Oke-Ose, Ilorin, Kwara State. The animals where kept in the animal house of the Faculty of Clinical Science, University of Ilorin, under light and dark cycle at room temperature. Proper aeration maintained by the use of wire gauze wooden cage. The rats were fed on growers’ feed from Bendel feeds Nigeria Limited, Ilorin, and distilled water throughout the duration of the experiment. The rats were arranged randomly into groups A-F, kept in different compartments of the cage. The rats were left under the above stated condition for two weeks so as to acclimatize with the condition of their new environment.

Grouping of animals
Group A- (n=4) control animals; these experimental animals were given pelleted feed and tap water ad libitum throughout the period of the experiment;
Group B- (n=4) experimental cryptorchidism was carried out this group;
Group C- (n=4) experimental cryptorchidism was carried out and then 100mg alpha tocopherol for 14days;
Group D- (n=4) animals were exposed to heat from hot oven for 28 days;
Group E- (n=4) animals were exposed to heat from hot oven for 28 days then 100mg/kg vitamin E for 14 days; group F- (n=4) animals were exposed to heat from light bulb for 28 days, group G- animals were exposed to heat from light bulb for 28 days then 100mg/kg vitamin E for 14 days.

Animal sacrifice and tissue collection
At the end of the substance administration, the rats were sacrificed 24hrs after the last day of the exposure using ketamine (0.2ml). The animals were pinned on the dissecting board. Dissection was done on the anterior abdominal wall (thoraco-abdominal sagittal incision) using the necessary surgical equipment/instruments. Blood was first collected via the apex of the heart with a hypodermic syringe, and the blood was centrifuged, the epididymis was also excised and placed in normal saline for semen analysis. The testis was removed and then fixed Bouin’s fluid.

Semen analysis
The testes from each rat were carefully exposed and removed. They were trimmed free of the epididymides and adjoining tissues. Epididymal sperm concentration: Spermatozoa in the right epididymis were counted by a modified method of Yokoi and Mayi, (2003). Briefly, the epididymis was minced with anatomic scissors in 5mL physiologic saline, placed in a rocker for 10 minutes, and allowed to incubate at room temperature for 2 minutes. After incubation, the supernatant fluid was diluted 1:100 with solution containing 5 g sodium bicarbonate and 1mL formalin (35%). Total sperm number was determined by using the new improved Neuber’s counting chamber (haemocytometer). Approximately 10µL of the diluted sperm suspension was transferred to each counting chamber of the haemocytometer and was allowed to stand for 5 minutes. This chamber was then placed under a binocular light microscope using an adjustable light source. The ruled part of the chamber was then focused and the number of spermatozoa counted in five 16-celled squares. The sperm concentration was the calculated multiplied by 5 and expressed as [X] x 106 /ml, where [X] is the number of spermatozoa in a 16-celled square (Oyewopo et al., 2010).
Sperm progressive motility: This was evaluated by an earlier method by Sonmez et al, (2005). The fluid obtained from the left cauda epididymis with a pipette was diluted to 0.5 mL with Tris buffer solution. A slide was placed on light microscope with heater table, an aliquot of this solution was on the slide, and percentage motility was evaluated visually at a magnification of x 400. Motility estimates were performed from three different fields in each sample. The mean of the three estimations was used as the final motility score. Samples for motility evaluation were stored at 350c. Sperm morphology: The sperm cells were evaluated with the aid of light microscope at x 400 magnification. Caudal sperm were taken from the original dilution for motility and diluted 1:20 with 10% neutral buffered formalin (Sigma-Aldrich, Oakville, ON, Canada). Five hundred sperm from the sample were scored for morphological abnormalities (Atessahin et al., 2006). Briefly, in wet preparations using phase-contrast optics, spermatozoa were
In this study a spermatozoon was considered abnormal morphologically if it had one or more of the following features: rudimentary tail, round head and detached head and was expressed as a percentage of morphologically normal sperm (Oyewopo et al., 2010).

Estimation of lipid peroxidation (Malondialdehyde)

Lipid peroxidation in the testicular tissue was estimated calorimetrically by thiobarbituric acid reactive substances TBARS method of Buege and Aust (1978). A principle component of TBARS being malondialdehyde (MDA), a product of lipid peroxidation. In brief, 0.1 ml of tissue homogenate (Tris-Hcl buffer, pH 7.5) was treated with 2 ml of (1:1:1 ratio) TBA-TCA-HCl reagent (thiobarbituric acid 0.37%, 0.25 N HCl and 1 5% TCA) and placed in water bath for 15 min, cooled. The absorbance of clear supernatant was measured against reference blank at 535nm. Concentration was calculated using the molar absorptivity of malondialdehyde which is 1.56 x 10^5 M^-1 cm^-1 and expressed as nmol/mg protein (Oyewopo et al., 2010).

Estimation of Glutathione level

Total GSH was estimated in various tissues by the method of Sedlak and Lindsay. Briefly, 5% tissue homogenates were prepared in 20 mM EDTA, pH 4.7, and 100 µl of the homogenate or pure GSH was added to 0.2 M Tris-EDTA buffer (1.0 ml, pH 8.2) and 20 mM EDTA, pH 4.7 (0.9 ml) followed by 20 µl of Ellman’s reagent (10 mmol/l DTNB in methanol). After 30 min of incubation at room temperature, absorbance was read at 412 nm in a Beckman DU-640 spectrophotometer. Samples were centrifuged before the absorbance of the supernatants was measured (Sedlak and Lindsay, 1968).

Estimation of Superoxide dismutase (SOD) level

Testicular tissues were transferred into 5ml ice-cold sucrose solution (0.25M) and homogenized. The homogenates were further centrifuged at 3000 rpm for 15 min to obtain the supernatant, which was then aspirated with Pasteur pipette into sample bottle, stored overnight at 4°C before being used for assays. Tissue activities of superoxide dismutase (SOD) were determined by the method of Marklund and Marklund, (1974).

Statistical analysis

Data collected were analysed using Microsoft Excel and one-way analysis of variance (ANOVA) followed by Tukey’s multiple comparison test (HSD) with the aid of SPSS V20. Data were presented as means ± SEM (standard error of mean) P value less than 0.05 (p<0.05) was considered statistically significant.

RESULTS

HISTOLOGICAL OBSERVATION

The micrograph from control group (group A) showed normal population of germinal epithelium, intact structure and arrangement of seminiferous tubules, presence of Leydig cell, and well vascularized. Micrograph from group B (cryptorchidic
animals) showed degeneration of spermatogenic cells and absence of spermatozoa in the lumen. Micrograph from Group C (cryptorchidic animals that were given alpha tocopherol) showed regeneration of spermatogenic cells and evidence of spermatogenesis due to presence of spermatozoa in the lumen. Micrograph from animals exposed to heat from hot oven (Group D), showed degeneration of basement membrane and clustering of spermatogonia cells, but Regeneration of of spermatogenic cells and normal spermatogenesis was evident in animals exposed to heat from hot oven followed by alpha-tocopherol (group E). Micrograph of animals exposed to heat from light bulb (Group D), showed distorted interstitium and cellular hyperplasia, which was reversed in animals that was given alpha-tocopherol.

**Figure (A-G):** photomicrographs showing the cyto-architecture of the testes across all experimental groups. Stain- Haematoxylin and Eosin (H & E). Mag- 400.

A- Control, B- cryptorchidism, C- crypt + vit E, D- hot oven, E- hot oven + vit E, F- light bulb, G- light bulb + vit E.

L-Lumen, White arrows- Spermatogenic cells, Black arrow- Basement membrane

Semen analysis
Table 1: percentage sperm count, sperm motility, life and ratio and sperm morphology across all groups.

Biochemical analysis
Table 2: malondialdehyde (MDA), glutathione (GLU) and superoxide dismutase (SOD) levels across all groups.

**DISCUSSION**

Cryptorchidism and hyperthermia caused desquamation and degeneration of germ cells which might be due to the destabilization of Sertoli cell membranes caused by lipoperoxidation, irregular shape of the seminiferous tubule, and degeneration of the interstitial tissues; this was in accordance to the report of Dutta et al., in 2013; Oyewopo and Togun in 2005 and Lee and Coughlin, in 2001. In cryptorchidic and hyperthermic animals exposed to alpha tocopherol (aT), lipoperoxidation of the membrane was impeded, conserving cell union and increasing the amount of germ cells in the seminiferous epithelium (Shikone et al., 1994).

The reduction in sperm parameters analysed as seen in figures 4.2 to 4.6 (i.e. sperm counts, motility, life and death ratio and morphology) in both cryptorchidic and heat treated animals might be due oxidative stress action that
cryptorchidism and hyperthermia cause to testicular tissues. Oyewopo & Togun in 2005 and Lee & Coughlin in 2001 also reported a significant reduction in sperm parameters of cryptorchidic and heat treated animals. But the effects of cryptorchidism and hyperthermia (heat) was reversed in animals treated with alpha tocopherol (Vitamin E) by not only returning sperm parameters to normalcy but also enhance the rate of spermatogenesis in the animals. My finding was corroborated by works of Saalu et al., in 2014 and Rosa et al., in 2011. Cryptorchidism and hyperthermia cause a decrease in antioxidant enzyme activity or an increase in the production of ROS, (i.e. superoxide anion, hydroxyl radical, nitric oxide and hydrogen peroxide) which stimulates lipoperoxidation which results in increase in malondialdehyde and reduced glutathione and superoxide dismutase levels in exposed animals. The work of Saalu et al., 2014; Rosa et al., 2011; Ishii et al., 2005 corroborate my findings. However administration of alpha tocopherol to the treated groups showed increase in cell’s endogenous antioxidant defence system, inhibiting ROS production and impede lipoperoxidation (as seen in figures 4.8 to 4.10). This result was corroborated by work of Saalu et al., 2014; Ayobami et al., 2013; Rosa et al., 2011 and Laila and Sahar in 2009.

CONCLUSION
Cryptorchidism and hyperthermia resulted in oxidative stress in exposed animals which is evident in distortion seen in the histological slides, reduced sperm parameters and antioxidant activities. This work showed that aT reduced apoptosis in germ cells, promoting cell survival and reducing histological alterations to the seminiferous epithelium.

REFERENCES
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